AΠ		
AD	 	

Award Number: DAMD17-01-1-0523

TITLE: UGT1A9 Genetic Polymorphisms and Raloxifene

Pharmacogenetics

PRINCIPAL INVESTIGATOR: Rebecca B. Raftogianis, Ph.D.

CONTRACTING ORGANIZATION: Fox Chase Cancer Center

Philadelphia, Pennsylvania 19111

REPORT DATE: May 2003

TYPE OF REPORT: Final

PREPARED FOR: U.S. Army Medical Research and Materiel Command

Fort Detrick, Maryland 21702-5012

DISTRIBUTION STATEMENT: Approved for Public Release;

Distribution Unlimited

The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.

# REPORT DOCUMENTATION PAGE

Form Approved OMB No. 074-0188

2

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing this collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to Washington Headquarters Services, Directorate for Information Operations and Reports, 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302, and to the Office of Management and Budget, Paperwork Reduction Project (0704-0188), Washington, DC 20503

1. AGENCY USE ONLY	2. REPORT DATE	3. REPORT TYPE AND	DATES COVERE	D
(Leave blank)	May 2003	Final (1 May 0	1 - 30 Apr	03)
4. TITLE AND SUBTITLE UGT1A9 Genetic Polymorph Pharmacogenetics  6. AUTHOR(S) Rebecca B. Raftogianis,			5. FUNDING N DAMD17-01-	
7. PERFORMING ORGANIZATION NAME Fox Chase Cancer Center Philadelphia, Pennsylvan E-Mail: RL_Blanchard@fccc	nia 19111		8. PERFORMIN REPORT NU	G ORGANIZATION MBER
9. SPONSORING / MONITORING				NG / MONITORING
AGENCY NAME(S) AND ADDRESS U.S. Army Medical Resear Fort Detrick, Maryland  11. SUPPLEMENTARY NOTES	ch and Materiel Comma	and	AGENCY H	EPORT NUMBER
12a. DISTRIBUTION / AVAILABILITY S				12b. DISTRIBUTION CODE
Approved for Public Rele	ease; Distribution Unl	imited.		
The goal of this DOD Breast Co in the human UDP-glucuronosyl enzyme expressed predominantl raloxifene (RAL). The pharmac associated with variable clinical to the known variation in RAL polymorphisms within the codin	ncept award was to identify a ltransferase gene, UGT1A9. It is the human liver, catalyze okinetics of RAL is known to efficacy. We hypothesized to harmacokinetics in humans.	We had previously des the glucuronidation be subject to significant that genetic variation. The aims of this pro-	letermined than and inactivaticant interinding in the human oposal were to	t UGT1A9, a metabolic tion of the antiestrogen vidual variation, possibly UGT1A9 gene contributed 1) identify genetic

14. SUBJECT TERMS  UDP-glucuronosyltransferase 1A9; pharmacogenetics; Raloxifene			15. NUMBER OF PAGES 7
			16. PRICE CODE
17. SECURITY CLASSIFICATION OF REPORT	18. SECURITY CLASSIFICATION OF THIS PAGE	19. SECURITY CLASSIFICATION OF ABSTRACT	20. LIMITATION OF ABSTRACT
Unclassified	Unclassified	Unclassified	Unlimited

allozymes with regard to capacity to glucuronidate RAL and 3) express variant UGT1A9 cDNAs in MCF-7 cells and

assess antiestrogenic response of cells to RAL.

## **Table of Contents**

Cover	1
SF 298	2
Introduction	4
Body	4
Key Research Accomplishments	5
Reportable Outcomes	7
Conclusions	7
References	N/A
Bibliography of Publications	N/A
List of Personnel	7
Appendices	N/A

#### INTRODUCTION

The goal of this DOD Breast Concept award was to identify and functionally characterize common genetic polymorphisms in the human UDP-glucuronosyltransferase gene, UGT1A9. We had previously determined that UGT1A9, a metabolic enzyme expressed predominantly in the human liver, catalyzes the glucuronidation and inactivation of the antiestrogen raloxifene (RAL). The pharmacokinetics of RAL is known to be subject to significant interindividual variation, possibly associated with variable clinical efficacy. We hypothesized that genetic variation in the human UGT1A9 gene contributed to the known variation in RAL pharmacokinetics in humans. The aims of this proposal were to 1) identify genetic polymorphisms within the coding regions of the human UGT1A9 gene, 2) functionally characterize recombinant UGT1A9 allozymes with regard to capacity to glucuronidate RAL and 3) express variant UGT1A9 cDNAs in MCF-7 cells and assess antiestrogenic response of cells to RAL.

### **BODY**

Aim 1. Identify common genetic polymorphisms in the human UGT1A9 gene. The UGT1A9 gene is part of a nested UGT1A gene family on human chromosome 2. The organization of this locus is such that alternative transcription initiation occurs at promoters of eight unique first exons, followed by splicing to common exons 2 through 5. Thus, eight unique UGT1A isoforms are expressed from this locus and those proteins differ in sequence at the amino terminal 530 amino acids by virtue of the unique exon 1 and they each share identical carboxy-terminal protein sequence that is encoded by the common exons 2 through 5. We and others have previously shown a lack of variation in gene sequence within the shared exons 2 through 5. Thus, genetic variation in UGT1A genes lies predominantly in the unique first exon. Therefore, we initially characterized common genetic variation in the UGT1A9-specific first exon.

Last year, we reported the identification of genetic polymorphisms within the 5'flanking region and 3' intron of UGT1A9. Table 1 describes those polymorphisms as well as their frequencies. Polymorphic loci were in genetic linkage such that different permutations of those polymorphisms defined eight apparent alleles. Of particular interest was the dT 9 or 10 variable length nucleotide repeat (VLNR) in the 5'flanking region of the gene. This position maps to the putative TATAA box of the UGT1A9 promoter.

Specific Aims 2 and 3. Functional Characterization of the UGT1A9 polymorphisms. Our original aims were to functionally characterize polymorphic UGT1A9 proteins (allozymes). However, none of the polymorphisms that we identified altered the encoded amino acid sequence of the protein. Therefore the experiments proposed would not be appropriate. Alternatively, we plan to evaluate the functional significance of the promoter polymorphism by comparing transcriptional activity of reporter constructs driven by the polymorphic promoter and by evaluating the correlation between level of

UGT1A9 transcript and genotype in cell culture systems. Progress on the latter aims has been hampered by a turnover in personnel and therefore an extension to this grant has been requested.

## KEY RESEARCH ACCOMPLISHMENTS

_	Identified five common genetic polymorphisms in the 5"flanking region and first intron of the human UGT1A9 gene.
_	Determined the frequency and linkage of each of those polymorphisms in a population of 65 healthy Caucasian Americans.
	Currently evaluating the functional significance of the VLNR in the putative promoter.

Table 1. Genetic Variation in the Human UGT1A9 Gene

UGT1A9 <u>Variable Loci</u>	Nucleotide Polymorphism	Frequency
	Т9	0.56
- 118 poly dT	T <sub>10</sub>	0.44
	G	0.98
- 87	A	0.02
	G	0.75
I1 152	A	0.25
	Т	0.59
I1 219	A	0.41
I1 313	С	0.56
11 313	A	0.44

<sup>-118</sup> and -87 refer to nucleotide positions upstream of the "A" in the ATG start codon. Il refers to intron 1 and the number following the "Il" designation refers to the nucleotide position downstream of the exon/intron junction.

## REPORTABLE OUTCOMES

Jeffrey Zalatoris, Ph.D. and Rebecca Blanchard Raftogianis, Ph.D., UDP-glucuronosyltransferase-specific glucuronidation inactivates 4-hydroxytamoxifen and raloxifene. Oral Presentation at 2002 DOD Breast Cancer Era of Hope Meeting in Orlando, Fla.

#### CONCLUSIONS

We set out to characterize common genetic polymorphisms in the human UGT1A9 gene. Surprisingly, no polymorphisms affecting encoded amino acid sequence were identified. However, five common polymorphisms were identified in 5'flanking and intron regions of the gene. Of particular interest is the variable length nucleotide repeat in the putative promoter region of the gene. We are currently testing the hypothesis that this polymorphism is of functional consequence. Genetically determined variation in UGT1A9 activity may be an important factor in the clinical response of individuals to drugs that are metabolized via this pathway. This study has contributed toward our knowledge of common genetic variation in the human UGT1A9 gene.

## REFERENCES

None

## **BIBLIOGRAPHY OF PUBLICATIONS**

None

#### LIST OF PERSONNEL PAID FROM GRANT

Stephen Beaupariant, Postdoctoral Associate Amanda Thistle, Scientific Technician